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IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Application of: Jon A. Wolff, )  
Vladimir S. Trubetskoy, Aaron G. Loomis, )  
Paul M. Slattum, Sean D. Monahan, )  
James E. Hagstrom, Vladimir G. Budker )  
Serial No.: 09/328,975 ) Examiner: Richard A. Schnizer  
Filed: 06/09/1999 )  
Group Art Unit: 1635 )

For: **Charge Reversal of Polyion Complexes**

Commissioner for Patents  
P.O. Box 1450  
Alexandria, VA 22313-1450

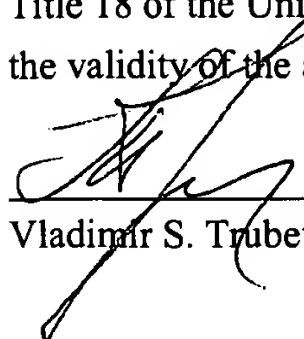
**DECLARATION UNDER 37 C.F.R. §1.131**

Dear Sir:

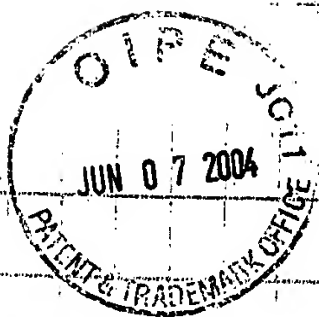
I, an inventor, Vladimir S. Trubetskoy, hereby declare as follows:

1. I am an inventor of the captioned application.
2. Photocopies of pages from my, Vladimir Trubetskoy's, personal laboratory notebook showing recharging of DNA/polycation particles beginning on December 16, 1997 accompany this Declaration.
3. It is known to me that the process performed in the notebook pages results in the formation of negatively charged tertiary complexes as described in the present specification.
4. The recharging process was conceived prior to the effective date of the Office Action prior art reference.
5. Developed of the recharging process occurred with due diligence from conception to the filing of the application.

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

  
Vladimir S. Trubetskoy      Date

6/7/04



1.5ul of react mix applied to

Albrecht preparative TLC silica plate and run in

product.

1.5ul of react mix applied to Albrecht preparative TLC silica plate and run in  $\text{CHCl}_3/\text{MeOH}$  (65:20) system.

Product band was scraped off the plate.

12/16/97

Work with  $\text{pDNA-hisMS}$

Part of silica (above) was washed with

(1)  $\text{CHCl}_3$

(2)  $\text{CHCl}_3/\text{MeOH}$  65:20

Substantial amounts of DSS (upper spot is present)

Whole amount of silica was washed with

(1)  $\text{CHCl}_3/\text{MeOH}$  65:10

(2)  $\text{CHCl}_3/\text{MeOH}$  65:30 → this fraction was evaporated

Work on recharging surface of caged DNA particles.

Caged particles are positively charged. If you add excess of polyanion it can recharge the surface to the opposite charge.

Caged particles were prepared in Bulker's conditions (p. 72)

After 2h of incubation of react mix at room to  
 The mixture was diluted twice with deionized  $H_2O$   
 and to 12% DNA/48% PLL capped, 500% of polymethac-  
 rylic acid (pMAA) were added.

No.	FI	Conc.
1	239.385	-10408 DNA/PLL (1:6) capped 1.7 DTBP
2	525.217	-22835 +500% pMAA
3	392.396	-17060 after centrif.
4	720.091	-31308 +150mM NaCl
5	481.248	-20923 after centrif.

Z-potential was also measured

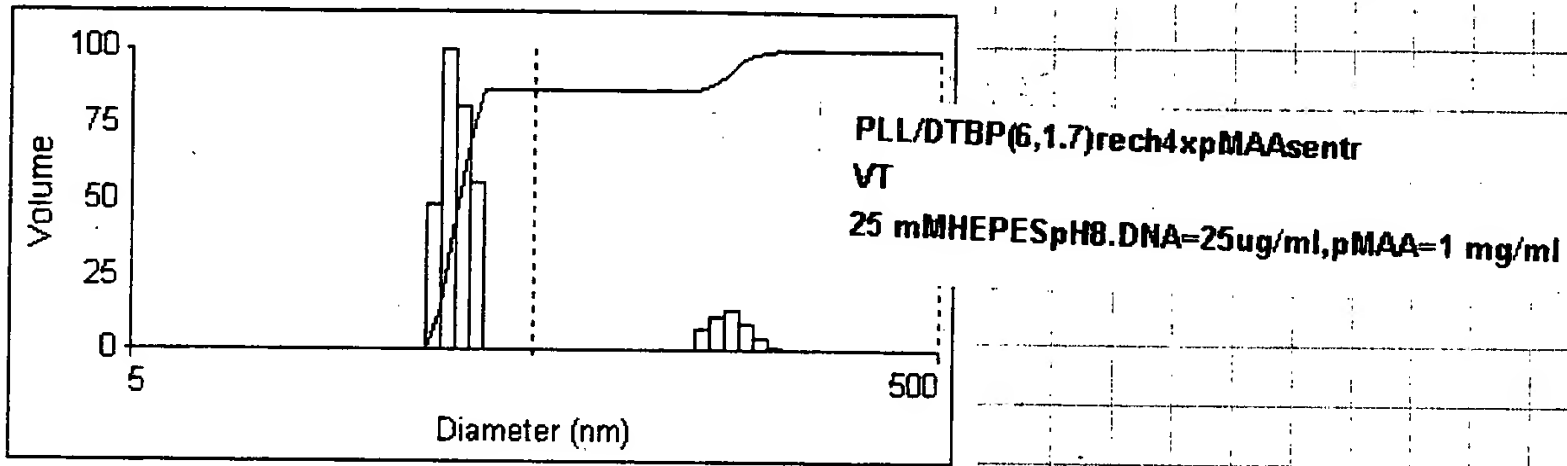
Run	Zeta Potential (mV)	Half Width (mV)
1	7.66	2.34
2	8.01	2.18
3	8.08	2.22
4	10.20	2.58
5	8.06	2.63
6	6.74	2.25
7	6.69	2.29
8	23.20	2.26
9	8.05	2.24
10	27.45	4.86
Mean	11.41	2.58
Std. Error	2.36	0.26

PLL/DTBP(6,1.7)nosalt (Run 10)  
 VT  
 ,DNA=17ug/ml, 17 mM HEPES, pH 8.0

Run	Zeta Potential (mV)	Half Width (mV)
1	-29.03	2.80
2	-7.70	4.06
3	-15.37	2.74
4	-25.43	3.53
5	-53.89	2.89
6	-16.53	2.89
7	-28.26	2.63
8	-24.13	3.00
9	-26.00	7.24
10	-35.16	4.16
Mean	-26.15	3.59
Std. Error	3.97	0.44

PLL/DTBP(6,1.7)+4xpMAAnosalt (Run 10)  
 VT  
 ,DNA=17ug/ml, 17 mM HEPES, pH 8.0

After addition of pMAA,  $I_{50}$  is increasing somewhat but still particle sizing



30/20/114 Ag/150

Basically the same effect was observed with dextran-sulfate(DS) as counterion. the mixture was as indicated on p 75 with exception that DS was added ~~is~~ instead of pMAA

Run	Zeta Potential (mV)	Half Width (mV)
1	33.22	2.41
2	27.98	2.61
3	20.17	3.26
4	26.99	2.22
5	10.37	2.35
6	27.01	2.06
7	33.33	2.24
8	25.83	4.46
9	28.83	2.93
10	29.39	2.18
Mean	26.31	2.67
Std. Error	2.13	0.23

PLL/DTBP(6,1.7)nosalt (Run 10)  
VT  
DNA=25ug/ml, 25 mM HEPES, pH 8.0, DS=0.5mg/ml

Run	Zeta Potential (mV)	Half Width (mV)
1	-7.34	2.32
2	-22.67	2.92
3	-13.63	2.19
4	-15.95	6.55
5	-2.55	3.97
6	-21.18	2.29
7	-25.78	2.10
8	-13.92	2.42
9	-11.06	2.01
10	-15.94	5.32
Mean	-15.00	3.21
Std. Error	2.23	0.50

PLL/DTBP(6,1.7)+500ugDSnosalt (Run 10)  
VT  
DNA=25ug/ml, 25 mM HEPES, pH 8.0, DS=0.5mg/ml

25 mg P-2636 Lot 75H5551

**SIGMA**  
POLY-L-LYSINE  
Hydrobromide (25988-63-0)

CAUTION: The chemical, physical and toxicological properties of this product have not been thoroughly investigated. Exercise due care.

Desiccate DP(vis) 251  
MW(vis) 52,400  
DP(LALLS) 252  
MW(LALLS) 52,700  
Store at less than 0°C M.W.(SEC-LALLS) 1.10  
For laboratory use only. Not for drug, household or other uses.  
MSDS available

SIGMA CHEMICAL CO. P.O. Box 14508 St. Louis, MO 63178 USA 314-771-5700

12/17/97 Titrations of DNA/PLL (1:6) caged and non-caged with dextran sulfate.

Beidker's solution was prepared as described in p. 72 this volume.

V = 1.5 ml (30  $\mu$ g DNA / 114  $\mu$ g PLL)

50  $\mu$ g - 500  $\mu$ g of dextran sulfate ( $M_w$  = 500 kDa, Sigma) were added to each sample and

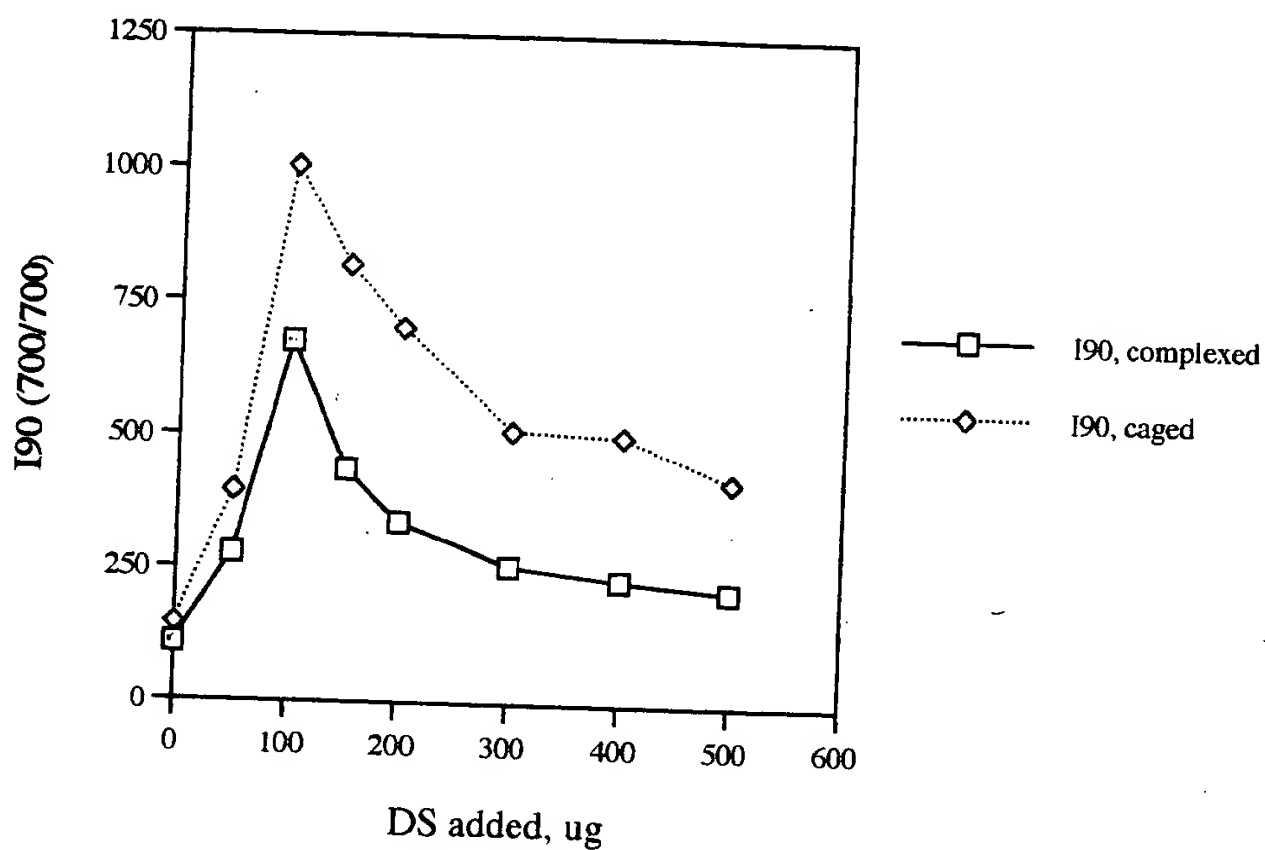
I<sub>50</sub>, F<sub>0</sub>TD, size and  $\zeta$ -potential were measured. Some non-caged samples were prepared in the same conditions.

TOTO concentrations: (8  $\mu$ l of stock TOTO into 20  $\mu$ l of 25 mM HEPES, pH 8.0; 10  $\mu$ l of sample  $\rightarrow$  0.5  $\mu$ l TOTO)

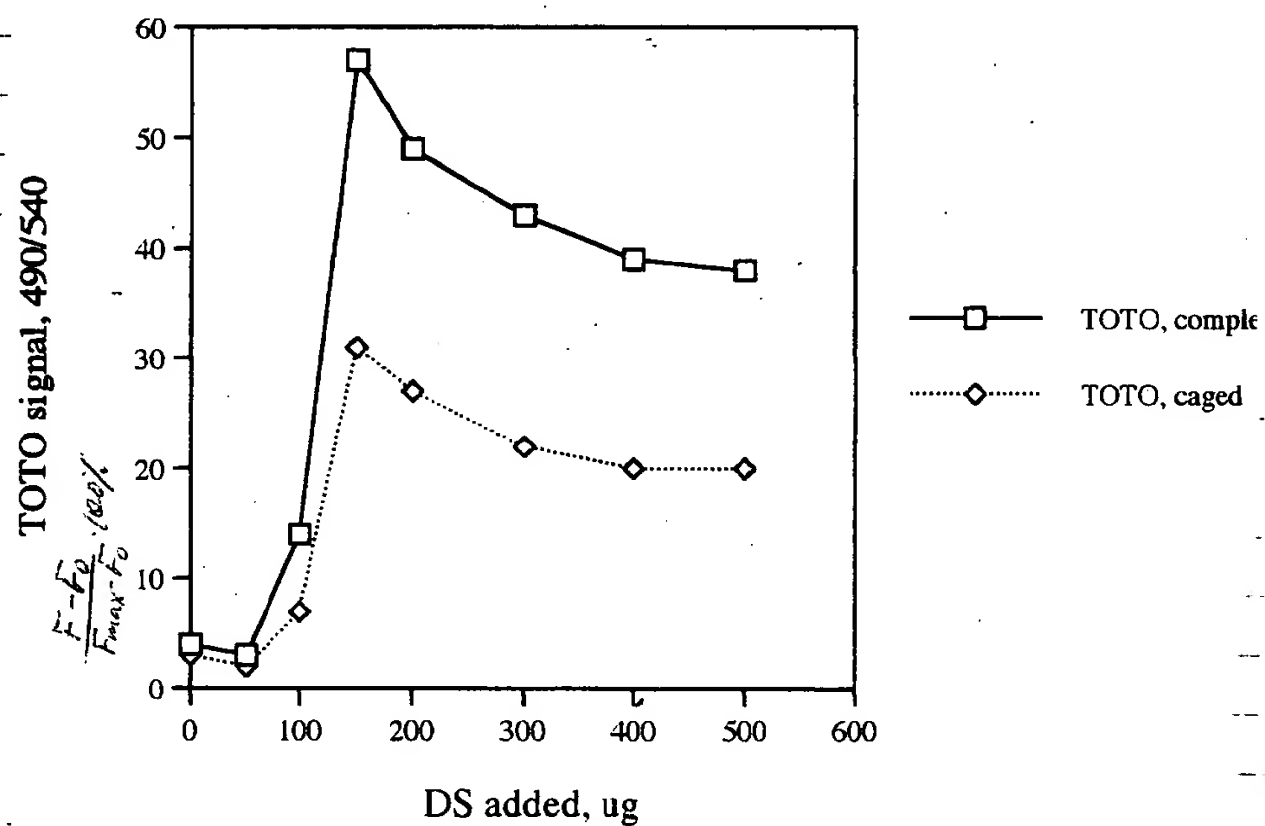
No. Caged	FI	Conc.	
		I <sub>50</sub>	200/300
1	145.344	-6319.3	0
2	395.046	-17175	50 $\mu$
3	1004.619	-43679	100 $\mu$
4	819.067	-35611	150
5	702.273	-30533	200
6	512.809	-22296	300
7	504.484	-21934	400
8	421.555	-18328	500 $\mu$
No.	FI	Conc.	TOTO
1	28.999	-1260.8	F <sub>0</sub>
2	687.116	-29874	F <sub>max</sub> 659
3	47.693	-2073.6	0
4	38.309	-1665.6	50
5	72.144	-3136.7	100
6	234.264	-10185	150
7	203.611	-8852.7	200
8	175.301	-7621.8	300
9	161.145	-7006.3	400
10	160.371	-6972.7	500

Complexed			
No.	FI	Conc.	
1	108.628	-4723.0	
2	278.651	-12115	
3	676.371	-29407	
4	435.570	-18937	
5	338.690	-14725	
6	258.092	-11221	
7	234.890	-10212	
8	215.716	-9379.0	
No.	FI	Conc.	
1	<del>96.057</del>	<del>4176.4</del>	
2	533.456	-23193	F <sub>max</sub> 490
3	64.342	-2797.5	0
4	60.599	-2634.7	50
5	111.724	-4857.6	100
6	322.742	-14032	150
7	284.332	-12362	200
8	253.480	-11020	300
9	236.314	-10274	400
10	230.641	-10027	500
11	43.587	-1895.1	F <sub>0</sub>

# Stabilization of DNA/PLL complexes (caged and complexed) with dextran sulfate



## Condensation of DNA/PLL/DS complexes (caged and complexed)

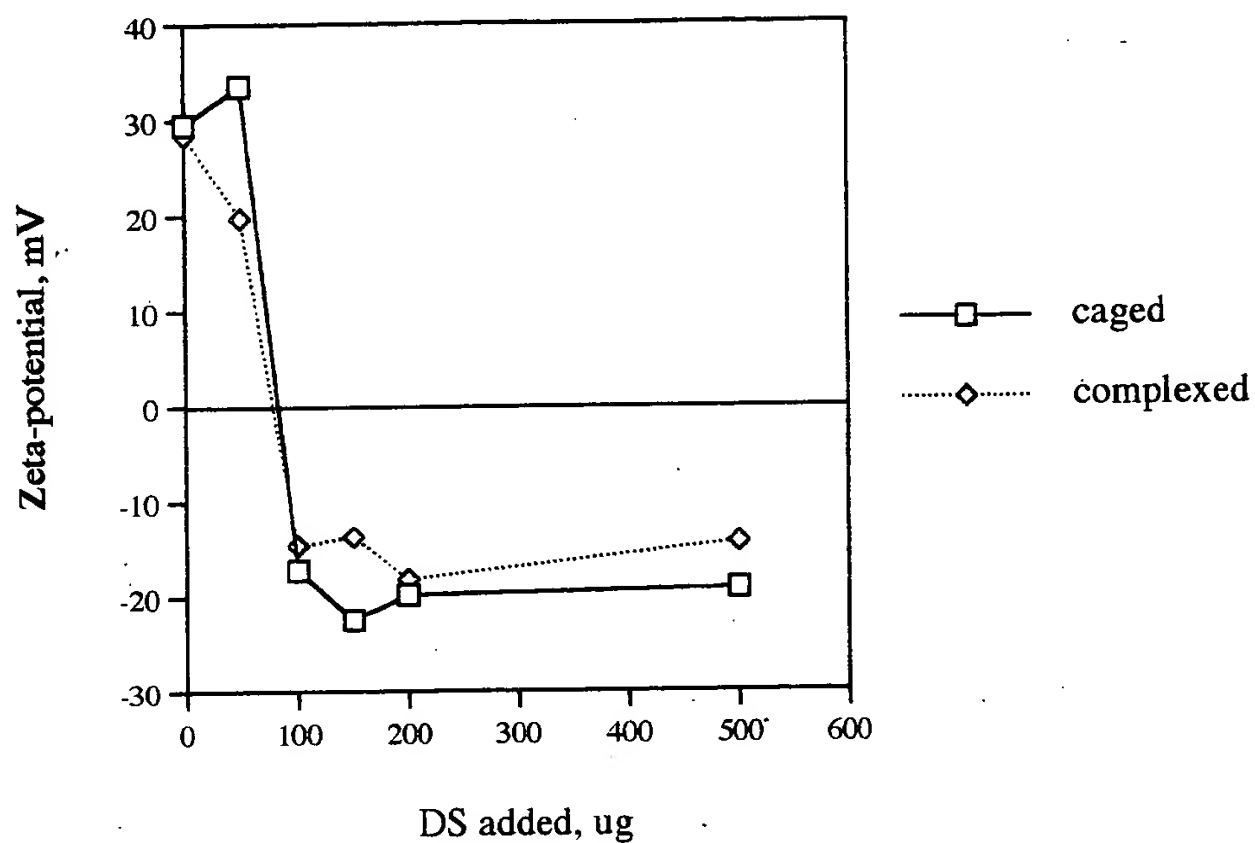


Complexed DNA/PLL were prepared in the same conditions as for caged but w/o  $\gamma$ -linking with DTBP.

$\zeta$ -potential is changed to opposite at 100 ug DS added.



## Zeta-potential of DNA/PLL/DS complexes, no salt



Run	Zeta Potential (mV)	Half Width (mV)
1	31.34	3.67
2	33.02	2.11
3	26.96	3.57
4	39.37	1.96
5	30.17	2.31
6	24.25	2.10
7	26.53	1.95
8	22.45	2.10
9	29.20	1.85
10	29.55	2.96
Mean	29.28	2.46
Std. Error	1.51	0.22

PLL/DTBP(6,1.7) (Run 10)  
VT  
DNA=20ug/ml, 25 mM HEPES, pH 8.0

Run	Zeta Potential (mV)	Half Width (mV)
1	49.36	3.06
2	44.33	1.83
3	37.00	1.80
4	33.83	3.36
5	39.11	2.34
6	27.81	1.81
7	28.67	4.53
8	11.79	1.82
9	36.92	1.84
10	28.00	3.19
Mean	33.68	2.56
Std. Error	3.30	0.30

PLL/DTBP(6,1.7)+50ugDS (Run 10)  
VT  
DNA=20ug/ml, 25 mM HEPES, pH 8.0

Run	Zeta Potential (mV)	Half Width (mV)
1	-18.29	1.86
2	-8.36	1.87
3	-6.31	1.93
4	-14.52	1.93
5	-14.56	1.89
6	-21.63	1.83
7	-18.70	1.81
8	-25.67	2.50
9	-22.83	2.45
10	-21.59	2.07
Mean	-17.25	2.01
Std. Error	1.99	0.08

PLL/DTBP(6,1.7)+100ugDS (Run 10)  
VT  
DNA=20ug/ml, 25 mM HEPES, pH 8.0



4	39.37	1.96	PLL/DTBP(6,1.7) (Run 10) VT ,DNA=20ug/ml, 25 mM HEPES, pH 8.0
5	30.17	2.31	
6	24.25	2.10	
7	26.53	1.95	
8	22.45	2.10	
9	29.20	1.85	
10	29.55	2.96	
Mean	29.28	2.46	
Std. Error	1.51	0.22	

Run	Zeta Potential (mV)	Half Width (mV)	PLL/DTBP(6,1.7)+50ugDS (Run 10) VT ,DNA=20ug/ml, 25 mM HEPES, pH 8.0
1	49.36	3.06	
2	44.33	1.83	
3	37.00	1.80	
4	33.83	3.36	
5	39.11	2.34	
6	27.81	1.81	
7	28.67	4.53	
8	11.79	1.82	
9	36.92	1.84	
10	28.00	3.19	
Mean	33.68	2.56	
Std. Error	3.30	0.30	

Run	Zeta Potential (mV)	Half Width (mV)	PLL/DTBP(6,1.7)+100ugDS (Run 10) VT ,DNA=20ug/ml, 25 mM HEPES, pH 8.0
1	-18.29	1.86	
2	-8.36	1.87	
3	-6.31	1.93	
4	-14.52	1.93	
5	-14.56	1.89	
6	-21.63	1.83	
7	-18.70	1.81	
8	-25.67	2.50	
9	-22.83	2.45	
10	-21.59	2.07	
Mean	-17.25	2.01	
Std. Error	1.99	0.08	

Run	Zeta Potential (mV)	Half Width (mV)	PLL/DTBP(6,1.7)+150ugDS (Run 10) VT ,DNA=20ug/ml, 25 mM HEPES, pH 8.0
1	-19.49	1.61	
2	-30.43	3.32	
3	-21.66	1.68	
4	-20.73	1.63	
5	-19.74	1.83	
6	-21.84	3.94	
7	-20.72	1.70	
8	-30.38	2.06	
9	-16.76	2.26	
10	-22.71	1.92	
Mean	-22.45	2.20	
Std. Error	1.42	0.25	

Run	Zeta Potential (mV)	Half Width (mV)	PLL/DTBP(6,1.7)+200ugDS (Run 10) VT ,DNA=20ug/ml, 25 mM HEPES, pH 8.0
1	-19.39	3.39	
2	-23.80	2.03	
3	-15.61	1.90	
4	-19.76	2.17	
5	-17.92	2.75	
6	-17.77	1.71	
7	-22.13	4.28	
8	-25.06	3.88	
9	-18.99	1.92	
10	-17.95	1.99	
Mean	-19.84	2.60	
Std. Error	0.93	0.29	

Run	Zeta Potential (mV)	Half Width (mV)	PLL/DTBP(6,1.7)+500ugDS (Run 10) VT ,DNA=20ug/ml, 25 mM HEPES, pH 8.0
1	-17.23	2.37	
2	-8.34	1.96	
3	-13.48	4.20	
4	-23.75	1.84	
5	-18.77	1.89	
6	-15.59	4.34	
7	-23.00	1.95	
8	-23.10	2.04	
9	-22.88	2.12	
10	-25.96	1.84	
Mean	-19.21	2.46	
Std. Error	1.76	0.31	

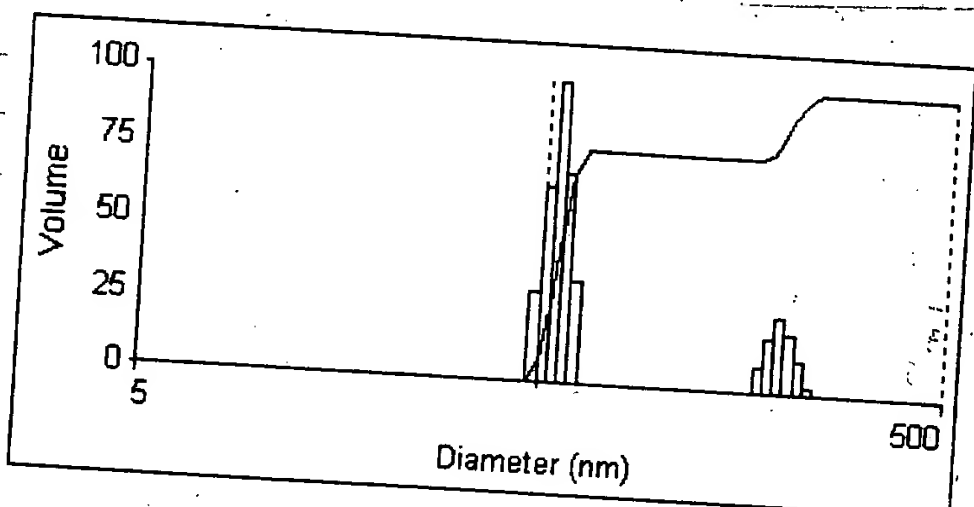
12/18/97

Work on recharged ~~ella~~ DNA colloid  
(precipitation in salt)

Samples prepared 12/17 (p77) were tested on precipitation  
upon addition of NaCl up to 150 mM.  
Was done with one sample 150  $\mu$ g DS - (close to neutrality point)

No.	$I_{50}$ (700/700) FI	Conc.	
1	396.680	-17246	compl 150 $\mu$ g DS
2	771.010	-33522	caged - " -
3	356.424	-15496	alt cent - " -
4	484.668	-21072	af. cent - " -
5	640.237	-27836	+ salt
6	667.412	-29017	+ salt
7	<del>618.884</del>	<del>26812</del>	
8	<del>681.295</del>	<del>29821</del>	
9	360.949	-15693	alt cent
10	400.766	-17424	alt cent

There is some  
aggregates formed  
after each step  
but significant  
amounts of particles  
stays in solution  
after addition of salt  
and centrifugation



Caged sample  
produced  
significant intens. 2  
(~1.2 Mcps) after  
centrifugation

DNA/PLL (1:6) caged stabilized w 150  $\mu$ g DS  
in 150 mM NaCl after centrifugation

12/23/97

Recharging the colloid with pMAA (not caged)

In standard settings: Complexes were formed  
at DNA = 50  $\mu$ g/ml in 25 mM HEPES pH 8, PLL/DNA = 6:1, V = 0.5 ml (25  $\mu$ g/55  $\mu$ l)  
→ then pMAA was added  
than each 0.5 ml was diluted to 1.5 ml with  
the same buffer

$I_{90}$ , TOTO and  $\zeta$  potential were measured

0 - 500  $\mu$  pMAA was added to each 25  $\mu$  DNA sample

No.	FI	Conc.
11	169.143	-7354.0

No.	FI	Conc.
1	177.670	-7724.8
2	695.225	-30227
3	995.999	-43304
4	320.682	-13942
5	603.757	-26250
6	316.927	-13779
7	456.850	-19863
8	305.441	-13280

No.	FI	Conc.
-----	----	-------

1	735.620	-31983
2	68.286	-2969.0
3	64.389	-2799.5
4	580.993	-25260
5	708.999	-30826
6	698.460	-30367
7	744.805	-32382
8	741.753	-32250
9	766.905	-33343
10	45.911	-1996.1

No.	FI	Conc.
-----	----	-------

1	125.019	-5435.6
2	339.144	-14745
3	1001.452	-43541
4	964.944	-41954
5	644.407	-28017
6	634.971	-27607

No.	FI	Conc.
-----	----	-------

1	393.592	-17112
2	54.019	-2348.7
3	45.936	-1997.2
4	47.624	-2070.6
5	359.647	-15636
6	225.945	-9823.7
7	206.946	-8997.7

32 - F<sub>0</sub>

pMAA  
Iso (500/600)

pMAA	mt DNA
690	0
23	25
19	50
535	100
663	200
653	300
699	400
696	500
721	

F<sub>0</sub>  
DS  
Iso

DNA	35S
0	16
25	7
50	9
100	321
200	187
300	168

47%

12/30/97

Recharging the DNA/PLL colloid (uncaged)

repetition of experiments from previous page

**TOTO**

No. FI Conc.

1	13.291	-577.87
2	894.401	-38886
3	94.502	-4108.8
4	339.541	-14762
5	844.788	-36729
6	901.778	-39207
7	931.606	-40504
8	961.974	-41824
9	978.774	-42555

**PMMA**

F <sub>0</sub>	881	100
F <sub>V</sub>	81	9.2
0	326	37.0
25	831	94.3
50	888	100.7
100	948	104.2
200	948	107.6
300	965	109.5

No. FI Conc.

1	14.718	-639.91
2	12.247	-532.48
3	11.329	-492.57
4	12.886	-560.26
5	12.353	-537.09
6	12.194	-530.17
7	12.591	-547.43

**PMMA - DNA**

No. FI Conc.

1	29.793	-1295.3
2	868.746	-37771
3	86.448	-3758.6
4	62.691	-2725.7
5	158.887	-6908.1
6	854.383	-37147
7	433.794	-18860
8	371.326	-16144
9	345.736	-15032

**DS**

F <sub>0</sub>	856	100
F <sub>V</sub>	74	8.6
0	50	5.8
25	146	17.0
50	842	98.4
100	421	49.2
200	359	41.9
300	333	38.9

No. FI Conc.

1	15.943	-693.17
2	12.170	-529.13
3	11.950	-519.57
4	12.479	-542.57
5	12.135	-527.61
6	14.364	-624.52
7	12.913	-561.43

**DS - DNA****I 90 (200/600)**

FI Conc. Z-potential

1	40.148	-6093.4	0
2	327.189	-14225	25
3	1008.335	-43840	50
4	753.784	-32773	100
5	559.717	-24335	
6	408.500	-17760	
7	332.728	-14466	

No.

FI Conc.

1	337.505	-14674	+
2	1008.335	-43840	+
3	1008.335	-43840	-
4	503.257	-21880	-
5	203.894	-8865.0	- 100
6	177.915	-7735.4	-
7	135.729	-5901.3	-

**PMMA**

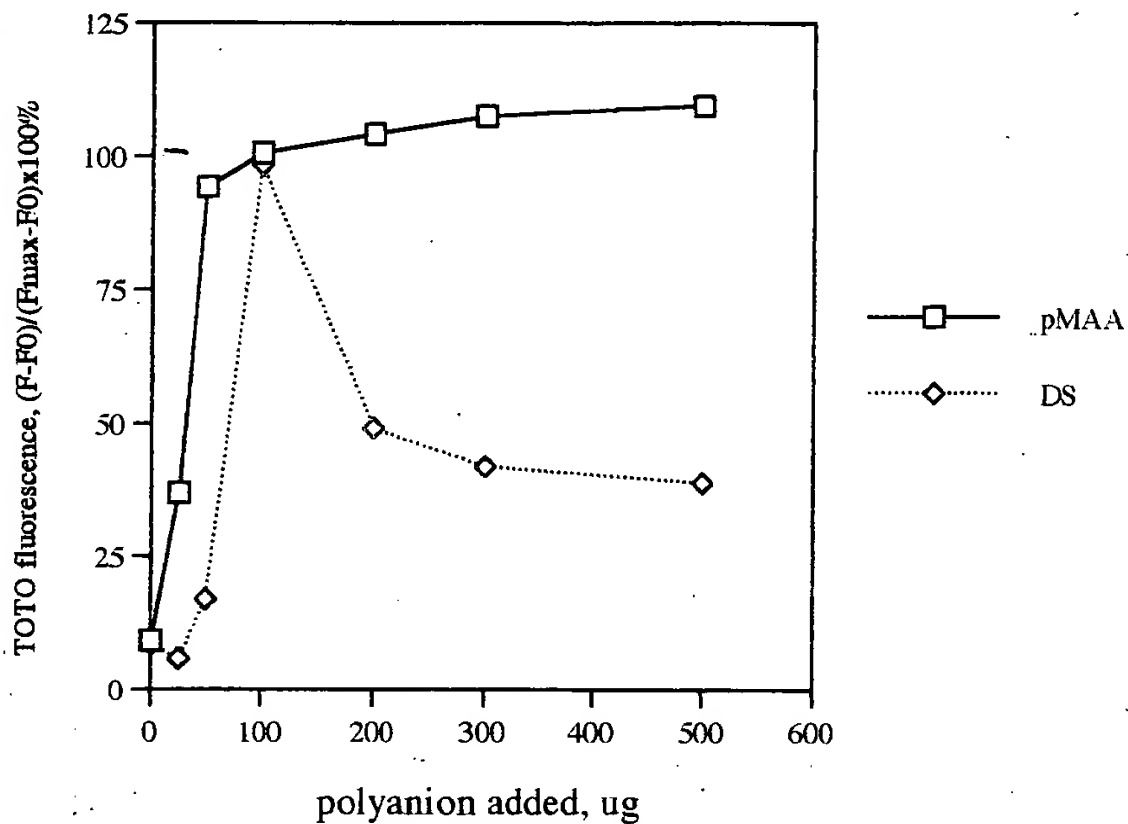
Conditions are the same as in p. 80.

TOTO signals from PMMA alone and

DS alone were measured.

polyanions did not change TOTO signals from DNA.

## Condensation of DNA/PLL(1:6) upon addition of polyanions



1/6/98

Precipitation of DNA/PLL complex after recharging with polyanion.

The complex DNA/PLL (1:6) + 200  $\mu$ g DS was prepared as in p. 80. 25  $\mu$ g / 95  $\mu$ g / 200  $\mu$ g in 0.5 ml 25 mM HEPES pH 8.0. then it was diluted up to 1.5 ml. 0.5 ml of this solution (17  $\mu$ g/ml) was tested for Iso

No.	FI	Conc.
1	173.291	-7534.4 DNA/PLL (1:6)
2 no	613.903	-26691 — " — +200 $\mu$ g DS
3 salt	219.280	-9533.9 DNA/PLL aft cent.
4	541.387	-23538 — " + 200 $\mu$ g DS + aft. cent.
5	723.397	-31452 DNA/PLL (1:6) in salt.
6 150mM	984.784	-42816 — " + 200 $\mu$ g DS in salt
7 salt	56.981	-2477.4 DNA/PLL in salt aft. cent.
8	588.275	-25577 — " + 200 $\mu$ g DS in salt aft. cent.